Protective activity of recombinant cytokines against Sendai virus and herpes simplex virus (HSV) infections in mice

Joji Iida, Ikuo Saiki, Chiaki Ishibara and Ichiro Azuma

The efficacy of recombinant cytokines such as marine interferoney (IFN-y), human gravulocyte colony-stimulating factor (C-SY), mouse gravulocytic-macrophage colony-stimulating factor (C-SY), motion gravulocytic-macrophage colony-stimulating factor (C-SY) and human interleukin-18 (IL-18) has been examined for augmentation of host resistance against Sendai wivas infection. IFN-y afforded protection when administrated intransauly has not intransauly surerul days before the infection. Intransaul administration of G-CSF one day before the infection was the most effective administration route and timing. Intransaul administration of Cotto SY and Intransaul administration of Cotto and the colony of the colony of the colony of the colony of the infection and in the colony of the colony of the colony of the internation on the same day as the infection. When each of the cytokines was administrate whose transaul administration of Cotto Cotto State of the cytokines was administrate whose transaction of the colony of the cytokines was administrate whose transaction of the cytokines was administrate whose transaction of the cytokines was administrate whose transaction of the cytokines was administrate whose whose transaction of the cytokines was administrate whose transaction of the cytokines was administrate whose the colony of the cytokines was administrate whose the colony of the cytokines was administrated whose the colony of the cytokines was administrated whose the colony of the cytokines of the cytokines.

Keywords: Sendai virus: cytokines; non-specific host resistance; infection; herpes simplex virus

Introduction

It has been found the sost resistance againg and herpes simples irus (HSV) infects and be enhanced by the admit ration of such bar till immunoadjuvants as whole probacterial popt saccharides (LPS) derived from Gr. negativ acterial nurars

dipeptide (MDP), which minimal unit immunoadjuvanticity⁻¹. Recently, host resistance to the infections has been shown to be augmented by murantipeptide, phosphatidylethanolamine (MTP-PE), which is a lipophilic derivative of MDP². We have previously reported that N-acatylmaranyl-analyl-siogaltaminyl-stepored that N-acatylmaranyl-analyl-siogaltaminyl-stepored that N-acatylmaranyl-analyl-siogaltaminyl-stepored that N-acatylmaranyl-analyl-siogaltaminyl-stepored that N-acatylmaranyl-analyl-siogaltaminyl-stepored that N-acatylmaranyl-analyl-siogaltaminyl-stepored that N-acatylmaranyl-analyl-siogaltaminyl-siogalta

MDP-Lys(L18) is a potent inducer of interleukin-IL-1) among macrophages in vito and colony-stimulating factor (CSF) in size³. Infections of Listeria mono-cytogenes, Salmonella typhinurulum and ectromelia virus, resistance to which largely depends upon macrophages, have caused the induction of CSF in sera³⁻¹. Alveolar macrophages exposed to influenza virus have produced IL-1 in virio-1. These results suggest that similar factors would be likely to have a role in natural host resistance to microbial infections. Recently, recombinant cytokines such as IL-1, granulocyte CSF (GM-CSF), granulocyte-mcrophage CSF (GM-CSF) as well as interferon-y

Institute of Immunological Science, Hokkaido University, Kita-15, Nishi-7, Kita-ku, Sapporo 060, Japan. (Received 1 November 1988) -y) have be four to prote mice against Listeria cytogenes, rudom is ar mosa Klebsiella partie. Escher codi, se process aureus, Serratia Candida alle si infections in normal or openic mice 13-13.

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this study, we attern id to evaluate the antiinfusious activity of G-CSF iM-CSF, IL-1 and IFN-y against Sendai virus infection in as well as against HSV infection in Cy-treated mice.

Materials and methods

Mice

Specific pathogen-free, male inbred Balb/c slc mice were obtained from the Shizuoka Experimental Animai Center and maintained in the Laboratory of Animai Experiment, Institute of Immunological Science, Hokkaido University, under laminar-flow conditions. All mice were used at the age of 4-5 weeks. Water and a pelleted diet (Nihon Nosan Kogyo Co. Ltd, Yokohama, Japan) were supplied ad libtum.

Reagent

Recombinant mouse interferon-9 (IFN-9) (Or, mg ml⁻¹; specific activity 10⁻¹ Um g⁻¹), prepared by Schering-Plough Corporation, was generously donated by the Suntory Co. Lid (Oska, Japan). Recombinant human granulocyte colony-stimulating factor (G-CSF) (50 g ml⁻¹), specific activity 3 × 10⁻¹ Um g⁻¹ vas described by Chugai Pharmaceutical Co. Lid (Toky, Japan). Recombinant human intertection-19 (IfV) (10, mm⁻¹) (10, mm⁻¹) Creaks Pharmaceutical Co. Lid (Toky, Japan). Recombinant human intertection-19 (IfV) (10, mm⁻¹) Creaks Pharmaceutical Co. Lid (Toky, Japan). Recombinant granulocyte-macrophase colony-

stimulating factor (GM-CSF) (1.17 mg ml⁻¹, specific activity 10° U mg⁻¹) was kindly provided by Sumitomo Pharmaceutical Co. Ltd (Osaka, Japan) Those recombinant cytokines were diluted with phosphate-buffered saline (PBS) containing 0.3% bovine serum albumin (BSA). In the perliminary experiment, we observed that the dilutent did not have any effect on protection against the dilutent did not have any effect on protection against or HSV infections. N²-Acetylmaramyl-Lysti (1.81) was donated by Daiichi Pharmaceutical Co. Ltd. Tokvo, Japan)

Protection avainst Sendai virus infection

Details of the methods have been reported previously6. The Sendai strain of parainfluenza type I virus was purchased from Flow Laboratories Inc., Rockville, MD. This virus was passaged for 11 generations in suckling C3H/He mice: after the 11th passage the lungs were homogenized in PBS and the supernatant fluid was dispersed in ampoules in 1 ml amounts, frozen and stored as the stock virus suspension at -70°C until use. For each experiment, an ampoule was thawed and 0,03 ml of the virus suspension was administered intranasally under light ketamine (Ketaral-50, Sankvo Co. Ltd. Tokyo, Japan) anaesthesia. When the infectious inoculum was assayed in LLC-MK2 cells (kindly donated by Dr H. Kida, Faculty of Veterinary Science, Hokkaido Univer-sity), it was found to contain one 10^{4.4} haemoadsorption (HAD) unit per 0.03 m re Were up to 21 days after th nfection. P alues wer calculated by apply the Mann-Whitne probability test to the mean su val times of the tre d group and up.

Protection against her infection

Herpes simplex virus (HSV) type I strain MacIntyre ATCC VR35's was provided by DrI. Sakaoka (School of Dentistry, Hokkaido University). A 10³⁻⁴ plaque-forming unit (pf.1, of HSV was injected intravenously in the mice which had received intrapertioneally cyclophosphamide (Sinongi Pharmacuttical Co. Ltd, Oaka, Japan) at a dose of 4 mg per mouse one day before the infection. Statistical analysis was performed as described above.

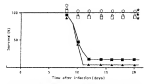


Figure 1. Protective activity of IFN-y against Sendai virus infection in mice. Seven or 8 Battly's nice were treated with various doses of IFN-y 3 days before infection with Sendai virus (10^{14} HAD per mouse). \bigcirc , 10^{14} U (10^{14} \square , \square U (10^{14}), \square , \square 10 U (10^{14}), \square 10 U (10^{14}), \square 2. \square 20 U (\square) 2.

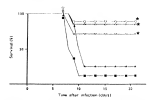


Figure 2. Effect of time intervals between treatment with IRN-y and infection with Sendai virus. Seven Balb'c mice were given 100 U IRN-y i.n. on various days before infection with Sendai virus (10° HAD per mouse). O, 5 days before, \(\triangle \), 3 days before; \(\triangle \), 3 days before; \(\triangle \), 4 days before; \(\triangle \), 5 days before; \(\triangle \), 2 days before; \(\triangle \), 5 days before; \(\triangle \), 2 days before; \(\triangle \), 1 day before; \(\triangle \), 9 \(\triangle \), 6 \(\triangle \), 9 \(\triangle \), 1 \(\triangle \), 9 \(\triangle \), 1 \(\triangle \), 9 \(\triangle \), 9 \(\triangle \), 1 \(\triangle \), 9 \(\triangle \), 1 \(\triangle \), 9 \(\triangle \), 1 \(\triangle \), 1 \(\triangle \), 9 \(\triangle \), 1 \(\triangle \), 1

Results

Protective activity of IFN-y on Sendai virus infection

The mice infected with Sendai virus died with severe pneumonitis 7-15 days after the infection (Figure 1). In the first experiment, we compared the protective activity adm stered intra ally (i.n.) or intrace that b eceived 10, 102 and J IFN-y i.n days re int on survived for 21 days 4-y i.v. was not effective infection. creas I (s). The su rate of the control group

e next examined the et of the timing of IFN-yac mistration on its prote we activity against Sendai view infection (Figure 2). Ameough i.n. administration of 10 IFN-y either 5 days, 3 days or 1 day before the infection, showed potent protective activity (88%, 88%, and 83%, respectively), postinfection administration (simultaneously with and Iday after the infection) had no effect (13%). The survival rate of the control group was 25%.

Protective activity of G-CSF and GM-CSF on Sendai virus infection

When the mice were given 2.0 µg G-CSF one day before infection, the survival rate was significantly higher than that of the control group (75%, Table 1, Experiment 1). Treatment 3 days before infection showed a slight protective activity (43%) and all the mice (which were injected simultaneously and I day after the infection) died within 21 days of infection. Subcutaneous (s.c.) or i.v. administration of 2.0 ug G-CSF 1 day before infection was not effective for protection agenst Sendai virus infection (Table 1. Experiment 2). Although the data are not shown, 2.0 μg G-CSF was the minimal effective dose for affording resistance to infection in our experimental conditions. The survival rate of the mice that received 2.0 µg G-CSF s.c. or i.v. I day before infection was similar to that of the control group (14%, 14% and 0%, respectively) (Table 1, Experiment 2). Intranasal administration of GM-CSF augmented host resistance to infection either 1 or 3 days before infection (Table 1, Experiment 3).

Protective activity of rIL-18 on Sendal virus infection

The results presented above show that i.n. administration of IFN-y and G-CSF before infection afforded a higher rate of protection against Sendai virus infection than i.v. or s.c. administration. Neither simultaneous

Table 1 Protective activity of G-CSF and GM-CSF against Sendai virus infection in mice

Experiment		Schedule*		Survivors/total		
no	Treatment	(day)	Route	on day 21	pb	
1	G-CSF	-1	i.n.	6/8	p < 0.901	
		~3	i.n.	3/7	0.02 < p < 0.05	
		0, +1	1. m.	0/7		
C	ontrol			0/7		
2	G-ÇSF	-1	i.n.	3/7	0.02	
		-1	S.C.	1/7		
		-1	i.v.	1/7		
Control				0/7		
3	GM-CSF	-3	l.n.	6/6	₽<0.001	
		-1	i.n.	6/6	p < 0.001	
Control				0/7		

⁴In each experiment, 2.0 µg G-CSF and GM-CSF were administered. Minus values indicate days before infection, plus values are days after infection. ⁵Probability values were calculated by the Mann-Whitney *U* test

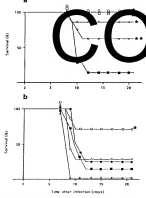


Table 2 Protective activity of G-CSF, and IFN-y against HSV infection in mice which received cyclophosphamide (Experiment 1)

	Trealment *			
Sample	On day	Daily dose	No. of survivors/total on day 21	p value
G-CSF	-4, -3, -2, -1	2,0 μα	1/7	
IFN-y	-4, -3 , -2 , -1	100 U	3/7	
MDP-Lys(L18)	-3, -1	100 µg	5/7	< 0 001

*Each of the samples was administered s.c. on the indicated days before infection. Cyclophosphamide was injected i.p. i day before infection at a dose of 4 mg. Mice were infected with HSV (10^{1,4} p.t.u.) Lv. *Probability values were calculated by the Mann-Whitney U lest





administration not postadministration of either cytokine was effective in increasing resistance to Sendai virus infection. As IL β is resistance to Sendai virus infection. As IL β is resistance to Sendai virus infection. As IL β is resistance to Sendai virus infection in the sendai virus infection by an infection in mice ^{3,14}, we examined its protective or therapeutic activity against Sendai virus infection by in. administration. Pretreatment with 0.2 μ g IL -1 β letther 3 days or 1 day before infection afforded protection against infection ($F | \mu \mu \sigma = 3.6$). Although the simultaneous administration of IL -1 β β had for infection) was remarkably effective, the post-administration of 0.2 μ g IL -1 β either 1 day or 3 days after infection was not effective.

Protective activity of IFN-7, G-CSF, GM-CSF and IL-1\(\beta\) on herpes simplex virus (HSV) infection in immunocompromised mice

Subcutaneous administration of MDP-Lys[L18], I day and 3 days before infection with HSV, almost completely protected the mice that had received intraperitoneally exclophosphamide one day before the infection (unpublished results). To evaluate the efficacy of IFN-y, G-CSF, GM-CSF and Li-fl gaginate HSV infection, various doses of the cytokines were administered subcutaneously four that MDP-Lys(L18) has a protective action, whereas

1FN-y, G-CSF and IL-1β were not effective. GM-CSF showed significant protective activity (Figure 4).

Discussion

We have already established a model of Sendui virus infection in mice, suitable for application to human pneumonitis caused by influenza virus infection, and was also reported that their, and aministration of synthetic adjuvants such as MDP derivatives and chitin derivatives and their new section of the s

The protection afforded by IFN-y, G-CSF and IL-18 against Sendai virus infection seemed to depend on the route or the timing of administration. Although i.n. administration was effective at a dose of 10 U, intravenous administration of 103 U was not effective (Figure 1). Intranasal administration of G-CSF 1 day before infection was also effective whereas i.v. or s.c. administration were not (Table 1, Experiment 2). Intranasal administration of G-CSF 1 day before infection was effective (Table 1. Experiment 1); i.n. on of G effective when it was, ninistere ther I or 3 da 1, Experiment 3 istration of these c before infection (To hese results suggest that i.n. adr kines is likely to cause an inflamn ory response, or the activates the immune system at e administration (lungs) and consequently stimul host resis ce a est the vira infection. Intranasal iniste of M was able to activate phagocytes in the lungs and the cells were able to suppress the growth of Sendai virus during the early phase of infection17. We observed that i.n. administration of IFN-y was more effective than i.v. administration in activating alveolar macrophages into their cytotoxic state against tumour cells (data not shown). Matsumoto et al. have reported that i.p. administration of G-CSF causes a significant increase in the peritoneal exudate cells and leads to the elimination of challenged hacteria in normal or in immunocompromised mice 15. CSF has been known to play a dual physiological role: it acts both to expand the macrophage-granulocyte population and also to enhance the functional activities of these cells 17. Subcutaneous administration of IL-1 Bleads, by its chemoattractive properties, within 1 h to the migration and accumulation of phagocytes around the administration site18. These findings suggest that the protection against Sendai virus infection afforded by intranasal administration of IFN-7, G-CSF, GM-CSF and 1L-1\beta may be attributable to the activation of alveolar macrophages or neutrophils. Only IL-1B showed any therapeutic activity against Sendai virus infection when it was administered simultaneously with the infection (2 h after infection) but it was not effective if it was administered either 1 day, 3 days or 5 days after infection (Figure 3a, b). The Sendai virus started to grow in the lung 10 h after infection and reached its maximum titre 2 days after infection⁷. Intranasal administration of IL-1β 2 h after infection may cause rapid accumulation of

phagocytes in the lungs before Sendai virus starts to grow. The precise details of the mode of action of $1L-1\beta$ are now being investigated.

It has been reported that macrophages have a major role in the protection of mice against HSV infection¹⁹⁻²¹. Of the four cytokines used in this study, only GM-CSF showed any significant protective activity against HSV infection in the Cy-treated mice.

In the present study we have attempted to evaluate the efficacy of IFN-y.G.CSF, GM-CSF and ILI-JB as protection against Sendai virus (local infection) and HSV (systematic infection) infections in normal and immunocompromised mice, respectively. The results show that the most effective route for the administration of these four cytokines against the Sendai virus infection is these four cytokines against the Sendai virus infection these four cytokines against the Sendai virus infection these four cytokines against the Sendai virus infection these results should be of clinical value in the protection of human patients against influenza and herres evirus infections.

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References

- Kirchner, H., Scott, M.T., Hirt, H.M. and Munk, K. Protection of mice against viral infection by Corynebacterium pervum and Bordefalla pertussis. J. Gen. Virol. 1978, 41, 97
- Spencer, T.C., Ganguly, R. and Waldman, R.H. Nonspecific protection of mice against influenza virus infection by local or systemic immunization with Bacillus Calmette-Guérin. J. Infect. Dis. 1977, 136, 171.
- 3 Gamgemi, J.D., Hightower, J.A., Jackson, R.A., Naher, M.H., Welsh, M.G. and Sige, M.M. Enhancement of natural resistance to influenza virus in lipopolysaccharide-rasponsiva and -nonreaponsive mice by *Propiophisecterium acres. Infact. Immun.* 1883, 39, 726
- Chedid, L., Parant, M., Audibert, F., Lefrancier, P., Choay, J. and Sala. M. Enhancement of certain biological activities of muramyl dipeptide derivatives after conjugation to a multipoly-jot-alaninol-poly(t-lysine) carrier. Proc. Natl Acad. Sci. USA. 1979, 76, 6557
- 5 Dietrion, F.M. Hockkoppel, H.K. and Lukas, B. Enhancomen of host resistance agents virus infections by MTP-E, a synthesic prophotic mutamy peptide. E Increased survival in mice and guines pigs after steple drug administration prior to infection, and the offect of MTPon instretion levels in seria and lungs Int. J. Immunopharmacol. 1986, 5, 931.
- 8 Ishihara, C., Hamada, N., Yamamoto, K., Ida, J. Azuma, I. and Yamamura, Y. Effect of muramyi dipophide and its stearoyi derivatives on resistance to Sendai virus infection in mice. Vaccine 1985, 3, 320.
- 7 Ishihara, C., Mizukoshi, N., Iida, J., Kato, K., Yamamoto, K. and Azuma, I. Suppression of Sendal virus growth by treatment with N°-acetylmuramyl--alaryl-tesiglutamine-N°-stearoyl--lysine in mice. Vector 1887, 8–265.
- 7a Ishihara, C., Iida, J., Mizukoshi, N., Yamamoto, N., Yamamoto, K., Kato, K. and Azuma, I. Effect of Nº-acetylmuramyl-u-alanyl-o-so-

- glutaminyl-M-stearoyl-c-lysine on resistance to herpes simplex virus type-1 infection in cyclophosphamide-treated mice. Vaccina, 1989. 7.
- in press 8 Saiki, I., Satio, S., Fujita, C., Ishida, H., Iida, J., Murata, J. et al. Induction of tumorcidal macrophages and production of cytokines by synthetic muramyl dipeptide analogues. Vaccina 1988, 6, 238
- Wing, E.J., Wahead, A. and Shadduck, R.K. Changes in serum colony-stimulating factor and monocytic properitor cells during Listeria monocytogenes infection in mice. Infact. Immun. 1984, 45, 180
 Trudgett, A., McNeill, T.A. and Killen, M. Granulcoyte-macrophage
- precursor cell and colony-stimulating factor responses of mice infected with Selmonella syphimurium. Infect. Immun. 1973, 8, 450 11 McNeill, T.A. and Killer, M. Hemopolietic colony forming cell responses in mice infected with extromelia virus. Infect. Immun. 1971, 4,
- 203 Roberts, N.J., Jr., Prill, A.H. and Mann, T.N. Interleukin 1 and interleukin 1 inhibitor production by human macrophages exposed to influenza virus or respiratory syncheli virus. Respiratory synchy virus is a potent inducer of inhibitor activity. J. Exp. Mod. 1966, 183,
- 511
 13 Ozaki, Y., Ohashi, T., Minami, A. and Nakamura, S. Enhanced resistance of mice to bacterial infection induced by recombinant human interlaukin-1*β*, Infect. Immun. 1987, 55, 1436
- human interlaukin-1β. Intect. Immun. 1987, 55, 1436
 Czuprynski, C.J. and Brown, J.F. Recombinant murine interleukin-

- 1β enhancement of non-specific antibacterial resistance Infect. Immun. 1987, 55, 2061
- Metamant, Levit, ad., 6001
 Metamano, M., Mastubera, S., Matsuno, T., Tamura, M., Hatton, K., Nomura, H. et al. Protective effect of human granulocyte colony-stimulating factor on microbial Infection in neutropenic mice. Infect. Immun. 1987, 55, 2715
- 16 lida, J., Une, T., Ishihara, C., Nishimura, K., Mizukoshi, N., Tokura, S. and Azuma, I. Stimulation of non-specific host resistance against Sendal virus and Escherichia coll infections by chitin derivatives in
- mice. Vaccine 1967, 5, 270

 17 Motcall, D. Acute antigen-induced elevation of serum colonystimulating factor (CSF) levels. *Immunology* 1971, 21, 427
- Stimulating labor (CSF) levels, immunology 911, 21, 401
 18 Oppenhelm, JJ, Kovacs, E.J., Matsushima, K. and Durun, S.K. Thera is more than one interleukin 1. Immunology Teday 1986, 7, 45
- There is more than one interleukin 1. Immunology Today 1986, 7, 45 19 Morahan, P.S., Glasgow, L.A., Crane, J.R. and Kern, E.R. Comparison of antiviral and antitumor activity of activated macrophages. Cell. Immunol, 1977, 28, 404
- Nirsch, M.S., Zisman, B. and Allison, A.C. Macrophages and age dependent resistance to herpes simplex virus in mice. J. Immunol. 1970, 104, 1160
- 1970, 104. 1160
 12 Zisman, B., Hirsch, M.S. and Allison, A.C. Salactive effects of anti-macrophaga serum, silica and anti-lymphocyte serum on pathogenesis of herpes virue infection of young adult mice. J. Immunol. 1970, 194, 1156

